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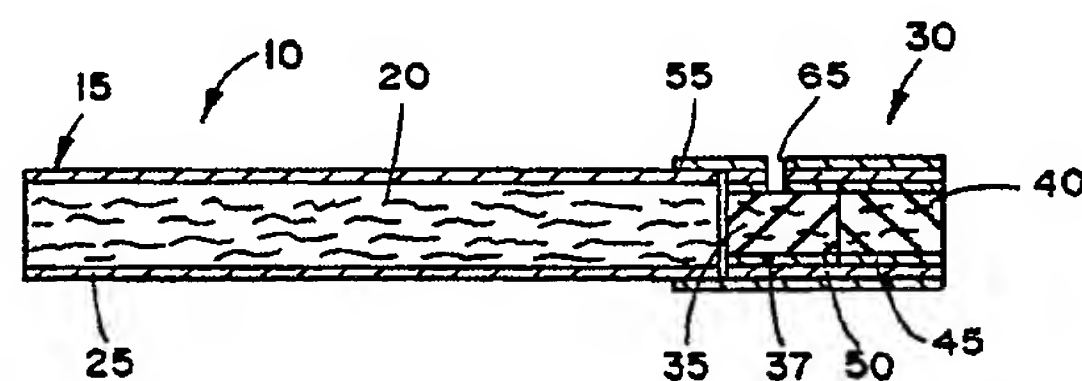
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**W-7000 Stuttgart 1(DE)**(54) **Cigarette and smokable filler material therefor.**

(57) A combustible smokable filler material includes an agglomerated matrix of an inorganic component (e.g., particles of calcium carbonate) and an organic component (e.g., an alginate). The smokable filler material also includes a binding agent, and an aerosol forming material. Tobacco extracts and/or pieces of tobacco laminae can be incorporated into the smokable filler material, and/or the smokable filler material can be blended with tobacco cut filler. Cigarettes are provided by wrapping the smokable filler material in a paper wrapping material. A typical paper wrapping material has a porosity of less than about 5 CORESTA units.

**FIG. 1****EP 0 419 975 A2**

## CIGARETTE AND SMOKABLE FILLER MATERIAL THEREFOR

### BACKGROUND OF THE INVENTION

The present invention relates to cigarettes and other smoking articles, and in particular to cigarettes, which when smoked, yield relatively low levels of incomplete combustion products, generate low amounts of sidestream "tar" and odor, and sustain smolder during FTC smoking conditions.

Popular smoking articles, such as cigarettes, have a substantially cylindrical rod shaped structure and include a roll or charge of smokable material, such as shredded tobacco material (e.g., in cut filler form), wrapped in a paper wrapper, thereby forming a so-called "smokable rod". Normally, a cigarette has a cylindrical filter element aligned in an end-to-end relationship with the smokable rod. Typically, a filter element includes cellulose acetate tow circumscribed by plug wrap, and is attached to the smokable rod using a circumscribing tipping material.

Typically, cigarettes are employed by the smoker by lighting one end thereof and burning the smokable rod. As such, smoke normally is provided by burning smokable material, which typically is tobacco cut filler. The smoker then receives mainstream smoke (e.g., mainstream tobacco smoke) into his/her mouth by drawing on the opposite end (e.g., the filter end) of the cigarette. As such, the smoker is provided with the pleasures of smoking (e.g., smoking taste, feel, satisfaction, and the like).

During the time that the cigarette is burning, sidestream smoke is generated. Sidestream smoke is smoke which directly enters the atmosphere from the lit end of the cigarette. Sidestream smoke diffuses into the atmosphere, and the characteristic visible nature and odor thereof may be perceived negatively by some individuals. The relative amount of visible sidestream smoke generated by a burning cigarette is related to the amount of sidestream "tar" generated by that burning cigarette. Typical commercially available cigarettes which burn tobacco cut filler, and have lengths of about 84 mm (e.g., having a smokable rod length of about 57 mm and a filter element length of about 27 mm), often yield about 25 to about 35 mg of sidestream "tar" per cigarette. See, Proctor et al, *Analyst*, Vol. 113, p. 1509 (1988), for an apparatus and technique for determining the sidestream "tar" of a cigarette.

Numerous cigarettes which reportedly yield relatively low levels of visible sidestream smoke have been proposed. See, for example, U.S. Patent Nos. 4,637,410 to Luke; 4,624,268 to Baker et al; 4,407,308 to Baker; 4,231,377 to Cline et al;

4,420,002 to Cline; 4,450,847 to Owens; 4,108,151 to Martin; 4,225,636 to Cline; 4,433,697 to Cline; 4,461,311 to Mathews et al; and 4,061,454 to Guess.

Through the years, there have been proposed various methods for altering the composition of mainstream tobacco smoke. For example, many tobacco substitute materials have been proposed, and a substantial listing of such materials can be found in U.S. Patent No. 4,079,742 to Rainer et al. In addition, tobacco substitute smoking materials having the tradenames Cytrel and NSM were introduced in Europe during the 1970's.

Numerous references have proposed articles which generate flavored vapor and/or visible aerosol. Most of such articles have employed a combustible fuel source to provide an aerosol and/or to heat an aerosol forming material. See, for example, the background art cited in U.S. Patent No. 4,714,082 to Banerjee et al.

Smoking articles which are capable of providing the pleasures associated with cigarette smoking, by heating but not necessarily burning tobacco, and without delivering considerable quantities of incomplete combustion products, are described in U.S. Patent Nos. 4,714,082 to Banerjee et al; 4,756,318 to Clearman et al; and 4,793,365 to Sensabaugh, Jr. et al. Such smoking articles employ a combustible fuel element for heat generation; and aerosol forming substances positioned physically separate from, and in a heat exchange relationship with, the fuel element. During use, heat generated by the fuel element acts to volatilize the aerosol forming substances, thereby providing an aerosol which resembles tobacco smoke. Such smoking articles yield extremely low levels of visible sidestream smoke as well as low levels of FTC "tar".

It would be desirable to provide a good tasting cigarette which provides good smoking satisfaction, provides relatively low mainstream gas phase yields, provides relatively low levels of incomplete combustion products, sustains smolder during FTC smoking conditions, and generates low levels of sidestream "tar" and hence low levels of visible sidestream smoke.

### SUMMARY OF THE INVENTION

The present invention relates to smoking articles incorporating tobacco in cut filler form and/or in a processed form. Preferred smoking articles have the form of a cigarette having two essential components: (i) a roll or charge of smokable ma-

terial, and (ii) an outer wrapping material (e.g., a paper wrapper) circumscribing the roll of smokable material. Cigarettes of the present invention incorporate a smokable filler material (described in greater detail hereinafter) as at least a portion of the smokable material thereof.

The preferred wrapping material, which surrounds the roll of smokable material to thereby form a "smokable rod", is a low air permeability cigarette paper wrapper. Highly preferred wrappers having a low air permeability or low porosity exhibit a porosity below about 5 CORESTA units. A CORESTA unit is a measure of the linear air velocity which passes through a 1 cm<sup>2</sup> area of wrapper at a constant pressure of 1 centibar. See CORESTA Publication ISO/TC 126/SC 1 N159E (1986).

One form of smokable material is a tobacco-containing smokable filler material. Such a smokable material of the present invention comprises an intimate mixture of (i) tobacco (e.g., shredded tobacco laminae, pieces of tobacco stems, milled tobacco laminae, tobacco fines, tobacco dust, or a tobacco extract or other form of processed tobacco), and (ii) an agglomerated matrix filler having, in intimate contact, an organic component and an inorganic component. Preferably, the agglomerated matrix filler is provided in particulate form. The organic component is an alginate or other organic material capable of providing an agglomerated matrix filler which is essentially insoluble in water at ambient conditions. Normally, the inorganic component is calcium carbonate. The smokable filler material includes a binding agent in intimate contact with the agglomerated matrix filler and tobacco. As such, the binding agent acts to maintain particles of the tobacco and agglomerated matrix filler together to form the smokable filler material. Such a tobacco-containing smokable filler material also can include certain flavoring agents (e.g., cocoa, licorice, organic acids, menthol, and the like) and/or aerosol forming materials (e.g., glycerin, propylene glycol, and the like) in intimate contact therewith. The tobacco-containing smokable filler material can be cast as a sheet from an aqueous slurry, provided as a sheet using a paper-making process, or provided in extruded form. Such a tobacco-containing smokable filler material can be employed individually as the sole smokable material of the cigarette, or that tobacco-containing smokable filler material can be physically mixed with (i.e., blended) or otherwise employed with other smokable materials, such as tobacco cut filler.

Another form of smokable filler material of the present invention comprises an intimate mixture of agglomerated matrix filler having, in intimate contact, an organic component and an inorganic component. Preferably, the agglomerated matrix filler is

provided in particulate form. The organic component is an alginate or other organic material capable of providing an agglomerated inorganic filler material which is essentially insoluble in water at ambient conditions. Normally, the inorganic component is calcium carbonate. The smokable filler material includes a binding agent in intimate contact with the agglomerated matrix filler. As such, the binding agent acts to maintain particles of agglomerated matrix filler together to form the smokable filler material. Such a smokable filler material also can include certain flavoring agents and/or aerosol forming materials in intimate contact therewith. The smokable filler material can be cast as a sheet from an aqueous slurry, provided as a sheet using a paper-making process, or provided in extruded form. Such a smokable filler material can be physically mixed with or otherwise employed with tobacco-containing smokable materials and/or tobacco cut filler.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1 through 4 are longitudinal sectional views of smoking articles of the present invention.

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

A preferred embodiment of a cigarette and smokable filler material of the present invention is shown in Figure 1. The cigarette 10 includes a generally cylindrical rod 15. The rod includes a roll of smokable material 20 wrapped in at least one layer of circumscribing outer wrapping material 25 (e.g., paper). The rod 15 is hereinafter referred to as a "smokable rod". The ends of the smokable rod 15 are open to expose the smokable material which is to be burned. The smokable rod is used by lighting one end thereof, and aerosol (e.g., smoke) is provided as a result of the combustion of the burning smokable material. As such, the smokable rod burns from the lit end thereof towards the opposite end thereof.

The cigarette 10 also includes a filter element 30 positioned adjacent one end of the smokable rod 15 such that the filter element and smokable rod are axially aligned in an end-to-end relationship, preferably abutting one another. Filter element 30 has a generally cylindrical shape, and the diameter thereof is essentially equal to the diameter of the smokable rod. The ends of the filter element are open to permit the passage of air and smoke therethrough. The preferred filter element has at least two filter segments. As shown in Figure 1, a first filter segment is positioned adjacent the



smokable rod, and preferably includes a carbonaceous filter material 35 circumscribed by a wrapping material 37; while a second filter segment is positioned at the extreme mouthend of the cigarette, and preferably includes a filter material 40, such as a gathered non-woven polypropylene web or cellulose acetate tow, circumscribed by a wrapping material 45. The filter material 40 of the segment preferably is a material which provides an aesthetically pleasing, white appearance. Each of the filter segments is manufactured using known filter rod making machinery. The two segments are combined using known plug tube combining techniques, and are held together using circumscribing wrap 50 so as to form the filter element.

The filter element 30 normally is attached to the smokable rod 15 by tipping material 55, which circumscribes both the entire length of the filter element and an adjacent region of the smokable rod. The inner surface of the tipping material 55 is fixedly secured to the outer surface of the plug wrap 50 and the outer surface of the wrapping material 25 of the smokable rod, using a suitable adhesive. The cigarette 10 can be manufactured using known cigarette making techniques and equipment. Optionally, a ventilated or air diluted cigarette is provided with an air dilution means such as a series of perforations 65 which extend through the tipping material 55, plug wrap 50 and wrapping material 37. Such ventilation can be provided to the cigarette using known techniques, such as laser perforation techniques.

Another preferred embodiment of a cigarette and smokable filler material of the present invention is shown in Figure 2. The cigarette 10 is generally similar to the cigarette described with reference to Figure 1, except that the smokable material has the form of a blend which is provided in a segmented fashion. At one end of the smokable rod 15 (i.e., at the end of the cigarette to be lit) is located a first segment 70 of smokable material. At the other end of the smokable rod 15 (i.e., at the end of the smokable rod adjacent the filter element) is located a second segment 75 of smokable material. Each segment is defined or identified in terms of its composition (i.e., the composition of each segment is different). The segments are aligned in an abutting, end-to-end relationship; however, there can be a certain amount of intermixing of smokable materials in the region where the two segments meet. The length which each segment of smokable material extends along the smokable rod can vary. However, the relative longitudinal length of the first segment relative to the second segment normally ranges from about 1:2 to about 2:1, with about 1:1 being preferred. Such smokable rods can be manufactured using apparatus described in U.S. Patent Nos. 4,009,722 to

Wahle et al and 4,516,585 to Pinkham.

For preferred cigarettes of the type shown in Figure 2, the first segment 70 normally includes of tobacco in some type of smokable form. Such a form of tobacco includes tobacco cut filler (e.g., tobacco laminae, processed tobacco materials, volume expanded tobacco filler, reconstituted tobacco filler materials, and the like, and blends thereof), and blends thereof with other smokable materials. Examples of processed tobacco materials are deproteinated reconstituted tobacco materials described in U.S. Patent Nos. 4,887,618 to Bernasek et al and 4,941,484 to Clapp et al, which are incorporated herein by reference. Another example of a processed tobacco material is a tobacco material processed according to the methods set forth in U.S. Patent Application Serial No. 484,587, filed February 23, 1990, which is incorporated herein by reference. Preferred cigarettes also have a second segment 75 which includes a smokable material or blend of smokable materials different in overall composition from the overall composition of the smokable material(s) of the first segment 70. The first segment 70 and/or the second segment 75 include at least one smokable filler material of the present invention.

Another preferred embodiment of a cigarette of the present invention is shown in Figure 3. The cigarette 10 is generally similar to the cigarette described with reference to Figures 1 and 2, except that the smokable material 20, which incorporates smokable filler material of the present invention is wrapped or contained in a processed tobacco sheet 80, or other inner wrapper material. The processed tobacco sheet 80 normally is a reconstituted tobacco sheet which is manufactured using a paper-making process, and a single layer of the sheet circumscribes the smokable material 20. The smokable material 20 wrapped in the processed tobacco sheet 80 is in turn wrapped in a single layer of circumscribing outer wrapping material 25 (e.g., cigarette paper).

Another preferred embodiment of a cigarette of the present invention is shown in Figure 4. The cigarette 10 is generally similar to the cigarette described with reference to Figures 1, 2 and 3, except that the filter element includes three segments. Segment 85, positioned between first filter material 35 and filter material 40, preferably is composed of a particulate matter such as activated carbon granules, magnesium silicate granules, silica gel particles, or the like.

The smokable material employed in the manufacture of the smokable rod can vary, and most preferably has the form of cut filler. As used herein, the term "cut filler" in referring to smokable materials is meant to include smokable materials which have a form suitable for use in the manufacture of

smokable rods for cigarettes. As such, cut filler can include smokable materials which are blended and are in a form ready for cigarette manufacture. Smokable materials normally are employed in the form of strands or shreds as is common in cigarette manufacture. For example, cut filler can be employed in the form of strands or shreds cut from sheet-like or "strip" materials. Such strip materials are cut into widths ranging from about 1/5 inch to about 1/60 inch, preferably from about 1/25 inch to about 1/35 inch. Generally, the resulting strands or shreds have lengths which range from about 0.25 inch to about 3 inches. Cut filler also can have an extruded form (e.g., extruded strands) or other physically processed form.

The smokable rods of cigarettes of the present invention include smokable filler material of the present invention. The smokable filler material can be employed in cut filler form.

One preferred type of smokable filler material of the present invention comprises an agglomerated matrix filler including an agglomerated matrix of an organic component and an inorganic component in intimate contact. The agglomerated matrix filler is in turn intimately mixed with a binding agent to provide the smokable filler material. Such a smokable filler material most preferably includes as part of the ultimate mixture, at least one aerosol forming material and/or at least one flavoring agent. If desired, other agents, which have the ability to alter the composition of the aerosol generated by the smokable filler material, can be incorporated into that smokable filler material. The agglomerated matrix filler normally includes about 2 to about 99, preferably about 5 to about 20 weight parts inorganic component for each weight part of organic component. Such a smokable filler material normally includes about 60 to about 90, preferably about 65 to about 85 weight percent agglomerated matrix filler; up to about 20, preferably about 2 to about 10 weight percent binding agent; up to about 20, preferably about 3 to about 15 weight percent aerosol forming material; and sufficient amounts of flavoring agent to provide the desired flavor characteristics. If desired, a carbonaceous material (e.g., pyrolyzed alpha cellulose) can be incorporated into the smokable filler material, usually in amounts of up to about 10, and sometimes up to about 30 weight percent, based on the total dry weight of the smokable filler material. However, such further carbonaceous material is not a necessary component of the smokable filler material, and the smokable filler material can be absent of such carbonaceous material. Such a smokable filler material is combustible and can be employed with (e.g., blended with) another smokable material (e.g., tobacco cut filler) in order to provide a cigarette of the present invention.

One preferred type of tobacco-containing smokable filler material of the present invention comprises an agglomerated matrix filler including an agglomerated matrix of an organic component and an inorganic component, which agglomerated matrix filler is in turn intimately mixed with some form of tobacco and binding agent, to provide smokable filler material. Such a tobacco-containing smokable filler material preferably includes as part of the ultimate mixture, at least one aerosol forming material and/or at least one flavoring agent. If desired, other agents, which have the ability to alter the composition of the aerosol generated by the smokable filler material, can be incorporated into that smokable filler material. The agglomerated matrix filler normally includes about 2 to about 99, preferably about 5 to about 20 weight parts inorganic component for each weight part of organic component. Such a tobacco-containing smokable filler material normally includes up to about 75, typically up to about 50 weight percent of some form of tobacco; up to about 80, preferably about 20 to about 60 weight percent agglomerated matrix filler; up to about 20, preferably about 2 to about 10 weight percent binding agent; up to about 20, preferably about 3 to about 15 weight percent aerosol forming material; and sufficient amounts of flavoring agent to provide desired flavor characteristics. If desired, a carbonaceous material (e.g., pyrolyzed alpha cellulose) can be incorporated into the smokable filler material, usually in amounts of up to about 10, and sometimes up to about 30 weight percent, based on the total dry weight of the smokable filler material. However, such further carbonaceous material is not a necessary component of the smokable filler material, and the smokable filler material can be absent of such carbonaceous material. Such a smokable filler material is combustible and can be employed individually as the sole smokable material of a cigarette of the present invention.

Another preferred type of smokable filler material of the present invention comprises an agglomerated matrix filler of an organic component and an inorganic component. The agglomerated matrix filler is intimately mixed with a binding agent and at least one aerosol forming material to provide the smokable filler material. Such a smokable filler material most preferably includes as part of the ultimate mixture, at least one flavoring agent and some form of tobacco. The agglomerated matrix filler normally includes about 2 to about 99 weight parts inorganic component for each part of organic component. Such a smokable filler material normally includes up to about 20, preferably about 3 to about 15 weight percent binding agent; greater than about 20, preferably about 25 to about 80, more preferably about 30 to about 50 weight per-

cent aerosol forming material; and less than about 80, preferably about 30 to about 70 weight percent filler component including the previously described agglomerated matrix filler. In particular, the filler component can include (i) all of the previously described agglomerated matrix filler, or (ii) a blend of the previously described agglomerated matrix filler with an inorganic filler material (e.g., precipitated calcium carbonate) and/or an organic filler material (e.g., tobacco). Amounts of flavoring agent sufficient to provide the desired flavor characteristics to the smokable filler material can be incorporated into the smokable material. If desired, a carbonaceous material (e.g., pyrolyzed alpha cellulose) can be incorporated into the smokable material, usually in amounts of up to about 10, and sometimes up to about 30 weight percent, based on the total dry weight of the smokable material. However, such further carbonaceous material is not a necessary component of the smokable material, and the smokable material can be absent of such carbonaceous material. The smokable filler material is combustible and can be blended with other smokable materials.

The tobacco-containing smokable filler materials of the present invention have some form of tobacco incorporated therein during its manufacture. The tobacco which is employed to provide such a tobacco-containing smokable filler material can have a variety of forms, including tobacco extracts, tobacco fines or dust, shredded or comminuted tobacco laminae, tobacco stems, volume expanded tobacco filler and other forms of processed tobacco, and the like, and combinations thereof. Tobacco extracts are processed forms of tobacco and are provided by extracting a tobacco material using a solvent such as water, carbon dioxide, a hydrocarbon, or a halocarbon, as well as various other organic and inorganic solvents. Tobacco extracts can include spray dried extracts; freeze dried extracts; heat treated extracts, such as those extracts described in U.S. Patent Application Serial Nos. 511,158, filed April 19, 1990 and 452,175, filed December 18, 1989; tobacco essences, such as those essences described in European Patent Application No. 326,370; and aroma oils and extracts described in U.S. Patent No. 4,506,682 to Mueller and U.S. Patent Application Serial No. 310,413, filed February 13, 1989.

The smokable filler materials of the present invention incorporate a binding agent. Examples of suitable binding agents include hydroxypropylcellulose such as Klucel H from Aqualon Co.; hydroxypropylmethylcellulose such as Methocel K4MS from The Dow Chemical Co.; hydroxyethylcellulose such as Natrosol 250 MRCS from Aqualon Co.; microcrystalline cellulose such as Avicel from FMC; methylcellulose such as Methocel A4M from The

Dow Chemical Co.; and sodium carboxymethylcellulose such as CMC 7HF and CMC 7H4F from Hercules Inc. Especially preferred binding agents include the alginates, such as ammonium alginate, sodium alginate, propylene glycol alginate and potassium alginate. The alginates, and particularly the high viscosity alginates, can be employed in conjunction with controlled levels of free calcium ions. Other binding agents include starches (e.g., corn starch), carrageenan, guar gum, locust bean gum, pectins and xanthan gum. Combinations or blends of binding agents (e.g., a mixture of guar gum and locust bean gum) can be employed.

The smokable filler materials of the present invention can have at least one aerosol forming material and/or at least one flavoring agent incorporated therein. The preferred aerosol forming materials include polyols (e.g., glycerin, propylene glycol or triethylene glycol), and any other materials which yield a visible aerosol, or combinations thereof. The aerosol forming material can be provided as a portion of the binding agent (e.g., when the binding agent is propylene glycol alginate). Combinations of aerosol forming materials can be employed. The flavoring agents can vary, and include menthol, vanillin, citric acid, malic acid, cocoa, licorice, and the like, as well as combinations thereof. See, Leffingwell et al, Tobacco Flavoring for Smoking Products (1972).

It is sometimes desirable to incorporate a caramelizing material into the smokable filler materials of the present invention. Caramelizing materials can act to improve (i) the integrity of the ash and fire cone of the cigarette, (ii) the appearance of the smokable filler material, and (iii) the flavor characteristics of the mainstream smoke of the cigarette. The caramelizing material can be incorporated into the smokable filler material during the preparation of that material and/or applied to the surface of that material (e.g., as a powder) after the manufacture thereof. Normally, the amount of caramelizing material which is employed to treat a particular smokable filler material is such that the resulting material which incorporates the caramelizing material includes up to about 20 weight parts of caramelizing material and greater than about 80 weight parts of the smokable material which is treated. Examples of suitable caramelizing materials include sugars, such as glucose, fructose and sucrose; and compositions such as Carob Powder Code 1739 from M. F. Neal, Inc.

The smokable filler materials of the present invention can be surface treated with certain substances. For example, the smokable filler materials can have powdered substances applied to the surface thereof. Exemplary substances include cocoa powder, licorice powder, powdered inorganic materials (e.g., potassium carbonate or iron oxide),



tobacco dust, finely divided tobacco laminae, or the like, or blends thereof. The surface treatment of the smokable filler materials can provide to those materials improved color and appearance, improved as characteristics, and improved flavor characteristics.

The agglomerated matrix filler has an inorganic component and an organic component. For example, calcium carbonate agglomerated using an alginate has an inorganic component and an organic component. Other inorganic components include calcium sulfate, magnesium oxide and magnesium carbonate. Ammonium alginate is an especially preferred alginate. Such an agglomerated matrix filler can be provided by preparing an aqueous slurry of calcium carbonate particles and hydrated alginate, and drying the slurry to form an agglomerated matrix of calcium carbonate (i.e., a matrix of a plurality of calcium carbonate particles spaced within a continuous or semi-continuous phase of alginate). If desired, the slurry can be volume expanded by incorporating a foaming agent therein. Examples of suitable foaming agents include linear sodium benzene sulfonates, linear alkyl sulfates and linear alkyl ethoxy sulfates. Preferably, a high solids content aqueous slurry of calcium carbonate and alginate is spray dried to provide agglomerated particles (e.g., normally spherical particles) of calcium carbonate particles and alginate. Alternatively, the slurry can be dried by the application of heat to provide a solid mass of agglomerated calcium carbonate and alginate, and the solid mass can be ground to yield particles of the desired size. Preferably, the amount of calcium carbonate relative to alginate ranges from about 99:1 to about 2:1, preferably about 20:1 to about 5:1, on a dry weight basis. Typically, the particles of calcium carbonate agglomerated using alginate are essentially insoluble in water under ambient conditions. In particular, the essentially insoluble character of the alginate in the agglomerated matrix filler tends to limit to a great degree any propensity of the agglomerated matrix filler to lose its agglomerated character when contacted with water under ambient conditions. The agglomerated matrix filler is rendered insoluble due to the interaction of the alginate with calcium ions of the calcium carbonate. If desired, the agglomerated matrix filler can be treated with a dilute solution of acid to decompose a portion of the calcium carbonate and the resulting calcium ions can act to render insoluble the alginate component of the agglomerated matrix filler.

The agglomerated matrix filler having an inorganic component and an organic component can incorporate a variety of other organic components. For example, the organic component can be pectin, which has a tendency to become essentially water insoluble upon interaction with calcium ions. Alter-

natively, agglomerated matrix filler having a polysaccharide organic component can be treated with divalent ions (e.g., calcium, barium, cobalt, iron or manganese ions) or trivalent ions (e.g., iron or aluminum ions) to render the polysaccharide essentially water insoluble. As yet another example, a slurry of polysaccharide material (e.g., ethylcellulose) and inorganic component particles can be provided in a non-aqueous solvent (e.g., alcohol) and dried, resulting in the formation of an agglomerated matrix filler which is essentially water insoluble.

Other inorganic materials can be incorporated as fillers in the smokable filler materials of the present invention. Such inorganic materials often have a fibrous, flake, crystalline, amorphous, hollow or particulate form. Examples of inorganic filler materials include calcium carbonate, calcium sulfate particles, magnesium oxide, magnesium hydroxide, perlite, synthetic mica, vermiculite, clays, thermally stable carbon fibers, zinc oxide, dawsonite, low density hollow spheres of calcium carbonate, glass spheres, glass bubbles, thermally stable carbon microspheres, calcium sulfate fibers, hollow ceramic microspheres, alumina, calcium carbonate particles agglomerated using a carbonaceous component, low density processed calcium carbonate, and the like.

The agglomerated matrix filler has the form of an agglomerated matrix of an inorganic component and a carbonaceous component. The inorganic component thereof can include particles of calcium carbonate, calcium sulfate, magnesium oxide, magnesium carbonate, and the like. A particularly preferred agglomerated matrix filler is agglomerated calcium carbonate, and most preferably, agglomerated precipitated calcium carbonate. Such an agglomerated matrix filler is prepared by providing an aqueous slurry of calcium carbonate particles and a binding material, and drying the slurry to form an agglomerated matrix of calcium carbonate (i.e., a matrix of a plurality of calcium carbonate particles spaced within a continuous or semi-continuous phase of binding material). Calcium carbonate particles which are employed to provide the agglomerated matrix typically exhibit a surface area of less than about 20 m<sup>2</sup>/g, frequently less than about 10 m<sup>2</sup>/g, and sometimes less than about 1 m<sup>2</sup>/g, as determined using the Brunauer, Emmett and Teller (BET) method described in J. Am. Chem. Soc., Vol. 60, p. 309 (1938). Typical binding materials are organic materials, such as cellulosic derivatives (e.g., sodium carboxymethylcellulose), and preferably are sugar containing materials, such as molasses, high fructose corn syrup, or Carob Powder Code 1739 from M. F. Neal, Inc. Other organic materials, such as pectins, alginates and the other previously described organic materials, also can be

employed.

Preferably, a high solids content aqueous slurry (e.g. about 40 to about 55 weight percent solids content slurry) of calcium carbonate and binding material is spray dried to provide agglomerated particles (e.g., normally spherical particles) of calcium carbonate particles and binding material. Alternatively, the slurry can be dried by the application of heat to provide a solid mass of agglomerated calcium carbonate and binding material, and the solid mass can be ground to yield particles of the desired size. Preferably, the amount of calcium carbonate relative to binding material ranges from about 20:1 to about 2:1, more preferably about 15:1 to about 4:1, on a dry weight basis. Normally, the inorganic particles agglomerated using saccharide and polysaccharide materials tend to lose their agglomerated character when contacted with water under ambient conditions, as a result of the propensity of the saccharide and polysaccharide materials to be soluble in water.

The agglomerated matrix of inorganic component and organic binding material is subjected to heat treatment. As such, volatile components from the organic binding material are expelled, and the organic binding material is calcined to form an essentially water insoluble, clean burning carbonaceous component. Normally, the heat treatment of the agglomerated matrix filler is provided under controlled atmosphere, in order to minimize or prevent oxidation of the binding material. See, for example, U.S. Patent Application Serial No. 414,833, filed September 29, 1989, which is incorporated herein by reference. Preferably, the heat treatment provides a binding material which is in the form of a carbonaceous material, and in turn, provides a means for agglomerating the particles of inorganic component into a matrix form. In particular, the particles of agglomerated calcium carbonate and binding material can be heat-treated (e.g., to a temperature of up to about 625° C, and usually up to about 600° C) using an oven, a fluidized bed, rotary calciner, belt calciner, or the like. For example, particles of spray dried calcium carbonate particles agglomerated using molasses can be heated in a fluidized bed having gaseous nitrogen flowing therethrough, heated at temperatures sufficient to heat the particles to about 300° C to about 625° C, and collected. The agglomerated matrix of inorganic component and organic binding material can be subjected to heat treatment sufficient to calcine the organic binding material by subjecting the agglomerated matrix to very high temperatures (e.g., up to about 900° C) for a short time period and under conditions sufficient to avoid decomposition of the inorganic component (e.g., when the inorganic component is calcium carbonate). However, if the inorganic component is calcium car-

bonate, and the calcium carbonate undergoes some decomposition during the calcining step, the agglomerated material can be re-carbonated by (i) exposing that material to carbon dioxide atmosphere, or (ii) dispersing that material in water and bubbling carbon dioxide into the dispersion.

After the calcining process, the agglomerated calcium carbonate particles normally have a calcium carbonate content of greater than about 80, frequently greater than about 90 weight percent and a carbon content of greater than about 3 weight percent. Normally, the resulting agglomerated particles are screened to sizes of about -50/+325 US Mesh, and often about -80/+200 U.S. Mesh. Preferred agglomerated calcium carbonate particles which have been calcined are essentially insoluble in water, are spherical in shape, are free flowing, and exhibit a bulk density of about 0.1 g/cm<sup>3</sup> to about 1.1 g/cm<sup>3</sup>, frequently about 0.3 g/cm<sup>3</sup> to about 1 g/cm<sup>3</sup>, using mercury intrusion techniques. As such, calcined agglomerated calcium carbonate particles provide an inorganic material having a bulk density less than about 2 g/cm<sup>3</sup>, and preferably less than about 1 g/cm<sup>3</sup>, which includes an inorganic component having a bulk density greater than about 2.5 g/cm<sup>3</sup>. Normally, such calcined agglomerated calcium carbonate particles exhibit a surface area of less than about 30 m<sup>2</sup>/g, and often about 10 m<sup>2</sup>/g to about 25 m<sup>2</sup>/g, as determined using the BET method.

If desired, the bulk density of the calcined agglomerated inorganic filler can be lowered by digesting away part of the inorganic component with an acidulant. For example, calcined agglomerated can be dispersed in water, and an aqueous hydrochloric acid solution can be added to the resulting slurry which is agitated. The acid reacts with the calcium carbonate, and essentially does not react with the carbonaceous component. Thus, the carbonaceous component acts to hold together the remaining calcium carbonate, while a portion of the calcium carbonate reacts to produce carbon dioxide gas and water soluble calcium chloride.

Another type of inorganic material which can be incorporated into smokable filler materials of the present invention is a low density inorganic filler. Such a filler is provided by providing particles of a calcium salt, decomposing the anion of the salt and contacting the particles with carbon dioxide. Examples of suitable salts include calcium propionate, succinate, tartrate, stearate, salicylate, palmitate, oleate, lactate, gluconate, citrate, ascorbate, acetylsalicylate and benzoate. Other suitable salts include calcium salts of saccharides and polysaccharides. Such salts are subjected to conditions sufficient to decompose the anion thereof, which usually involves subjecting the salt to heat treatment under carbon dioxide atmosphere.



One method for providing a low density inorganic filler involves heating calcium lactate particles screened to -80/+170 US Mesh at about 600°C for about 8 hours under a steady 228 ml/min. flow of carbon dioxide gas, so as to provide a material which has undergone about a 65 percent weight loss. About 20 weight parts of the material is charged into about 80 weight parts water and the resulting slurry is contacted with sufficient hydrochloric acid solution to lower the pH thereof to about 6.8. The material then is removed from the water, washed with water, dried, and screened to a particle size of -80/+170 US Mesh. Such material is greater than about 95 weight percent calcium carbonate, and exhibits a bulk density of about 0.4 g/cm<sup>3</sup>, as determined using mercury intrusion techniques.

Typically, the smokable filler materials of the present invention are provided by forming an aqueous slurry of binding agent and the other components of that smokable filler material, casting the slurry as a sheet, and drying the cast material to form a relatively dry, workable sheet. Techniques and equipment for casting a slurry as a sheet will be apparent to the skilled artisan. Other materials, such as calcium acetate, potassium carbonate, pH control agents, urea, amino acids, potassium chloride and/or calcium hydroxide, can be incorporated into the slurry. Sequestering agents (e.g., diammonium hydrogen orthophosphate, sodium citrate, potassium hexametaphosphate or tetrasodium pyrophosphate) can be incorporated into the slurry in amounts sufficient to control the free calcium ion concentration of the slurry. The cast material can be dried at ambient temperatures or at elevated temperatures. Further, an aqueous solution of calcium salts can be applied to the cast slurry. The resulting dried sheet can be cut or broken into "strip" form, and later can be cut or shredded into cut filler form. The smokable filler materials of the present invention can be provided using a paper-making process. In particular, an aqueous slurry of a cellulosic material (e.g., softwood pulp, hardwood pulp, flax fibers and/or shredded tobacco stems) and the previously described filler can be cast as a mat on a fibrous belt or wire screen, and dried to the desired moisture level. Normally, a slurry, dispersion or solution of flavoring agents, tobacco extracts, tobacco parts, aerosol forming materials, and the like, can be applied to the mat (e.g., as a spray), and the resulting mat can be dried further to form a sheet. The resulting dried sheet can be cut or broken in "strip" form, and later can be cut or shredded into cut filler form. Techniques and equipment for making a paper-type sheet will be apparent to the skilled artisan.

The smokable filler materials of the present invention can be extruded into the desired shape

using suitable extrusion techniques. See, for example, the types of processes described in U.S. patent No. 4,880,018 to Graves, Jr. et al, which is incorporated herein by reference. Alternatively, an aqueous slurry of the components of the smokable material and an alginate binding agent can be extruded into an aqueous solution of calcium ions (e.g., an aqueous solution of calcium chloride), collected and dried. If desired, extruded smokable filler materials can be physically processed (e.g., subjected to treatment using rollers, etc.) and formed into the desired shape.

The smokable rods of cigarettes of the present invention often include a physical mixture or blend of smokable materials. The blend can include two or more smokable filler materials of the present invention, or a physical mixture of at least one smokable filler material of the present invention with at least one other smokable material. Certain preferred cigarettes often include within such a blend, a sufficient amount of at least one of the smokable filler materials of the present invention such that the smokable material within each cigarette comprises at least about 1 percent of the carbonaceous material, based on the total weight of the blend. In particular, cigarettes having such types of smokable filler materials and having low porosity paper outer wrappers (e.g., having outer wrappers having less than about 5 CORESTA units) have the propensity to sustain smolder (e.g., not self-extinguish), when smoked under FTC smoking conditions. FTC smoking conditions consist of 35 ml puffs of 2 second duration, taken every 60 seconds.

The smokable filler materials of the present invention can be blended with tobacco cut filler. The type of tobacco can vary, and can include flue-cured, Burley, Maryland and Oriental tobaccos, as well as the rare and specialty tobaccos, and blends thereof. Such tobacco cut filler can be provided in the form of tobacco laminae; volume expanded or puffed tobacco laminae; processed tobacco stems such as cut-rolled or cut-puffed stems; reconstituted tobacco materials, such as (i) deproteinated tobacco materials described in U.S. Patent Nos. 4,887,618 to Bernasek et al and 4,941,484 to Clapp et al, (ii) a phosphate-containing reconstituted tobacco material described in U.S. Patent Nos. 3,353,541 and 3,420,241 to Hind et al, and 3,386,449 to Hind, as well as U.S. Patent Application Serial Nos. 406,637, filed September 13, 1989 and 461,216, filed January 5, 1990, (iii) a reconstituted tobacco material described in U.S. Patent Application Serial No. 272,156, filed November 16, 1988 and Tobacco Encyclopedia, edit. by Voges, p. 389, TJI (1984), (iv) the reconstituted tobacco materials described in U.S. Patent Application Serial Nos. 416,332, filed September 29, 1989 and

414,833, filed September 29, 1989; or blends thereof.

Smokable materials can be cased and top dressed as is conventional during various stages of cigarette manufacture. For example, flavoring agents can be applied to the smokable material as is commonly performed when cigarette cut filler is processed. Suitable flavoring agents include vanillin, cocoa, licorice, menthol, and the like. Flavor modifying agents can be applied to the smokable material. A flavor modifying agent in the form of levulinic acid can be applied to the smokable filler material (e.g., in amounts ranging from about 0.01 to about 2 percent, normally from about 0.1 to about 1 percent, preferably about 0.2 to about 0.6 percent, based on the dry weight of the smokable material). Another flavor modifying agent in the form of potassium carbonate can be applied to the smokable material (e.g., in amounts of less than about 5 percent, normally about 2 to about 3 percent, based on the dry weight of the smokable material). Aerosol forming materials and humectants, such as glycerin and propylene glycol, can be applied to the smokable material. Such components conveniently are applied to the smokable material as casing and top dressing components.

The preferred wrapping material which provides the smokable rod is a cigarette wrapping material having a low air permeability value. Such a wrapping material normally has an air permeability of less than about 5 CORESTA units, often less than about 3 CORESTA units, and frequently less than about 1 CORESTA unit. Typical wrapping materials are cigarette paper wrappers. Suitable wrapping materials are cigarette paper wrappers available as DD-71-1, DD-71-6, MTR-1021, P-2831-60-2, P-2831-60-3, P-2831-60-4, P-2831-60-5, P-2674-110, P-2831-60-1 and DD-100-2 from Kimberly-Clark Corp. Suitable low porosity cigarette paper wrappers are commercially available, and can have various levels of burn chemicals, fluxing agents, etc., incorporated therein. Particularly preferred are cigarette paper wrappers which include an amount of a polymeric film forming agent sufficient to provide a paper having the desirably low air permeability value. For example, a sufficient amount of a solution of a polymeric (e.g., carboxymethyl cellulose or ethylcellulose) film forming agent can be applied to a paper wrapper. The selection of the polymeric film forming agent will be apparent to the skilled artisan.

The optional polymeric film forming agent can be applied to the paper wrapper during the manufacture of the paper, or applied as a print or paint after manufacture of the paper is complete. Typically, the film forming agent is applied to the paper as a dilute solution (e.g., at a concentration of about 0.2 to about 5 weight percent relative to the

solvent) for ease of processing. The amount of film forming agent applied to the paper wrapper depends upon factors such as the permeability of the paper and the film forming capabilities of the film forming agent. Typically, the amount of film forming agents employed ranges from about 1 to about 10 percent, based on the dry weight of the paper. For example, a 5 weight percent solution of ethylcellulose in ethanol or sodium carboxymethylcellulose in water can be applied to cigarette paper using a size press, and the paper can be dried to provide a non-wetting, moisture resistant paper wrapper having a porosity of less than about 1 CORESTA unit, preferably less than about 0.5 CORESTA unit.

The smokable rods and the resulting cigarettes can be manufactured in any known configuration using known cigarette making techniques and equipment. Smokable rods often include smokable material wrapped in a single layer of wrapping material, although a double layer of two types of wrapping materials can be employed. See, for example, U.S. Patent Application Serial No. 528,302, filed May 24, 1990, which is incorporated herein by reference.

Cigarettes having smokable rods which are double wrapped with two layers of wrapping material preferably include one of the previously described low porosity paper wrappers as the outer wrappers. The inner wrapper can vary, but typically is a tobacco-containing wrapping material. Exemplary inner wrappers are paper wrappers which include about 3 parts Java tobacco stem parts and about 1 part wood pulp, and are available from Kimberly-Clark Corp. as p-2249-115 and P-2831-23-3. Other suitable inner wrapping materials include tobacco parts and carbonaceous materials, and are available from Kimberly-Clark Corp. as P-2540-94-A, P-2540-94-C and P-2540-94-D. The inner wrapping materials (i) can include burn chemicals (e.g., potassium citrate, potassium acetate or potassium succinate), and/or (ii) act as a substrate for flavors (e.g., menthol or vanillin) or flavor precursors (e.g., vanillin glucoside or ethylvanillin glucoside).

Typically, the smokable rod has a length which ranges from about 30 mm to about 70 mm, preferably about 35 to about 60 mm; and a circumference of about 17 mm to about 27 mm, preferably about 22 mm to about 25 mm. Short smokable rods (i.e., having lengths from about 30 to about 50 mm) can be employed, particularly when smokable materials having a relatively high packing density are employed.

The packing density of the smokable material contained within the outer wrapping material can vary. Typical packing densities for smokable rods of cigarettes of the present invention range from

about 150 to about 400 mg/cm<sup>3</sup>. Normally, packing densities of such smokable rods range from about 200 to about 380 mg/cm<sup>3</sup>, frequently about 250 to about 360 mg/cm<sup>3</sup>, particularly when relatively short (i.e., less than 50 mm long) smokable rods are employed.

The cigarettes of the present invention preferably include a filter element, and most preferably a filter element having more than one segment. For example, a preferred filter element has two or more filter segments. Typically, the segments of the preferred filter elements each have lengths which ranges from about 10 mm to about 30 mm; and circumferences of about 17 mm to about 27 mm, preferably about 22 mm to about 25 mm. The plug wrap which circumscribes the filter material of each filter segment typically is a conventional paper plug wrap, and can be either air permeable or essentially air impermeable.

Preferred filter materials of one of the filter segments include carbonaceous materials (e.g., activated carbon particles, charcoal particles, or carbon paper). An example of a particularly preferred filter material is provided by gathering a tobacco/carbon paper available as P-144-BAC from Kimberly-Clark Corp. Such filter materials reduce the levels of certain gas phase components from the mainstream smoke which passes to the mouth of the smoker. As such, preferred filter materials of that segment act to reduce the levels of any smoke components which may provide an off-taste or other undesirable characteristics to the mainstream smoke.

Preferred filter materials of another of the filter segments normally include fibrous materials. An example of a suitable filter material is a gathered nonwoven polypropylene web. A particularly preferred nonwoven polypropylene sheet-like web is available as PP-100-F from Kimberly-Clark Corp. Another example of a suitable filter material is a cellulose acetate tow. Particularly preferred cellulose acetate tow items include (i) 8 denier per filament/40,000 total denier, and (ii) 8 denier per filament/15,000 total denier, (iii) 8 denier per filament/25,000 total denier, and (iv) 8 denier per filament/30,000 total denier. Plasticizers, such as triacetin, propylene glycol or triethyl citrate, can be combined with the filler materials.

Another filter segment can have a filter material in the form of a gathered web of nonwoven thermoplastic (i.e., hydrophobic) fibers in intimate contact with a water soluble tobacco extract so as to provide an extract-containing filter material. A highly preferred web is a nonwoven web of polypropylene fibers available as PP 200 SD from Kimberly-Clark Corp. Exemplary filter segments and filter elements are described in U.S. Patent Application Serial Nos. 414,835, filed September 29, 1989 and 518,597,

filed May 3, 1990. Such segments can provide enhanced flavor characteristics to the mainstream smoke which passes therethrough.

Yet another filter segment can include a tobacco paper material as the filter material. For example, a filter material can have the form of a gathered web of tobacco paper available as P-144-B from Kimberly-Clark Corp.

The filter element segments suitable for use in this invention can be manufactured using known cigarette filter making techniques. Filter elements can be manufactured from cellulose acetate tow using known techniques. Filter elements can be manufactured from carbon paper, tobacco paper and a sheet-like nonwoven polypropylene web using filter making techniques described in U.S. Patent No. 4,807,809 to Pryor et al, which is incorporated herein by reference. Alternatively, particles of charcoal or activated carbon can be incorporated into the filter element using a so-called "triple filter" configuration by positioning the particles between two segments of suitable filter materials.

The filter elements can have low, moderate or high filtration efficiencies. Preferred filter elements have minimal mainstream aerosol (i.e., smoke) removal efficiencies while maintaining the desirable draw characteristics of the cigarette. Such minimal smoke removal efficiencies are provided by "low efficiency" filter elements. Low efficiency filter elements have a minimal ability to remove mainstream smoke particulates. See, Keith in Schemeltz's The Chemistry of Tobacco and Tobacco Smoke, p. 157 (1972). Generally, low efficiency filter elements provide less than about 40 weight percent mainstream smoke particulate removal efficiency.

Tipping material circumscribes the filter element and an adjacent region of the smokable rod such that the tipping material extends about 3 mm to about 6 mm along the length of the smokable rod. Typically, the tipping material is a conventional paper tipping material. Tipping materials of varying porosities can be employed. For example, the tipping material can be essentially air impermeable, air permeable, or treated (e.g., by mechanical or laser perforation techniques) so as to have a region of perforations, openings or vents thereby providing a means for providing air dilution to the cigarette. The total surface area of the perforations and the positioning of the perforations along the periphery of the cigarette can be varied in order to control the performance characteristics of the cigarette.

For air diluted or ventilated cigarettes of the present invention, the amount of air dilution can vary. Typically, the amount of air dilution for an air-diluted cigarette is greater than about 25 percent, and frequently greater than about 40 percent. The upper limit for air dilution for a cigarette typically is



less than about 75 percent, more frequently less than about 65 percent. As used herein, the term "air dilution" is the ratio (expressed as a percentage) of the volume of air drawn through the air dilution means to the total volume of air and aerosol (i.e., smoke) drawn through the cigarette and exiting the extreme mouthend portion of the cigarette. See, Selke et al, Beitr. Zur Tabak. In. , Vol. 4, p. 193 (1978).

Cigarettes of the present invention, when smoked, provide a flavorful mainstream aerosol. The mainstream aerosol of such cigarettes can yield low levels of incomplete combustion products as well as low levels of gas phase components. The cigarettes burn at an acceptable rate, and maintain static smolder, at least when smoked under FTC smoking conditions. The cigarettes, when smoked, have an ash and fire cone which is not overly cohesive, and hence, is not overly long. However, the cigarettes also provide an ash and fire cone which exhibit good integrity.

Cigarettes of the present invention, when smoked, generally yield less than about 20 mg, preferably less than about 10 mg of sidestream "tar" per cigarette, as determined using the apparatus and techniques described by Proctor et al, Analyst , Vol. 113, p. 1509 (1988). Such cigarettes normally provide more than about 5 puffs, preferably more than about 6 puffs per cigarette, when smoked under FTC conditions. Normally, cigarettes of the present invention provide less than about 20 puffs, and often less than about 15 puffs, when smoked under FTC conditions.

The following examples are provided in order to further illustrate the invention but should not be construed as limiting the scope thereof. Unless otherwise noted, all parts and percentages are by weight.

#### EXAMPLE 1

##### A. Preparation of a Tobacco-Containing Smokable Filler Material

A filler is provided as follows:

Into a high shear blender is charged about 500 parts tap water and about 5 parts of a high viscosity ammonium alginate available as Amoloid HV from Kelco Division of Merck & Co., Inc. The resulting mixture is agitated at ambient temperature so as to disperse the alginate in the water and hydrate the alginate. Then, about 95 parts of precipitated calcium carbonate available as USP Light from Pfizer Inc. is added to the water/alginate mixture. The resulting slurry is agitated at moderate rate for about 10 minutes until a consistent slurry is pro-

vided.

The slurry is spray dried using an Anhydro Type Lab S1 spray dryer having an air atomized nozzle at an inlet temperature of about 290 °C and an outlet temperature of about 220 °F. The spray dryer is equipped with a Masterflex Pump System using a Model 7016-20 head supplied by Cole-Parmer Instrument Co. The nozzle air pressure is 4 psi, and the liquid feed rate is that provided by setting the pump speed control to 2. The resulting spray dried particles have a generally spherical shape. The particles are screened to a particle size of -80/+325 U.S. Mesh.

The spray dried particles are free flowing, are essentially insoluble in water at ambient temperature, resist wetting, and remain in an agglomerated state when contacted with water under ambient conditions. The particles each are an agglomerated matrix of a plurality of precipitated calcium carbonate particles spaced within an alginate.

The smokable material is provided as follows:

Into a high shear blender containing 225 parts tap water and set at low speed is charged about 5 parts of a high viscosity ammonium alginate available as Amoloid HV from Kelco Division of Merck & Co., Inc. The resulting mixture is agitated at ambient temperature so as to disperse the alginate in the water and hydrate the alginate. Then, 20 parts glycerin and 25 parts tap water are charged into the water/alginate mixture. Then, 17.2 parts of the previously described agglomerated calcium carbonate is charged into the mixture. The resulting slurry is agitated until a consistent slurry is provided.

The slurry is cast to about a 0.025 inch thickness onto a high density polyethylene sheet and air dried. The resulting sheet light brown color and a flexible character. The sheet is cut at 32 cuts per inch using a paper shredder.

The shreds of sheet are dusted with about 8.5 parts of finely divided Pennsylvania cigar tobacco laminae.

##### B. Preparation of a Cigarette

Cigarettes substantially as shown in Figure 3 are provided as follows:

The cigarettes each have a length of about 84 mm and a circumference of about 24.8 mm, and include a smokable rod having a length of about 57 mm, a first filter segment having a length of about 15 mm and a second filter segment having a length of about 12 mm. The first and second filter segments form a filter element. Each filter segment is attached to each smokable rod using nonporous tipping paper. For each cigarette, the tipping paper circumscribes the filter element and about a 4 mm

length of the smokable rod in the region adjacent the filter element. The filter elements are ventilated to about 60 percent air dilution by providing a ring of perforations through the tipping paper and plug wrap of the filter element circumscribing the cigarette about 12 mm from the extreme mouthend thereof.

The smokable rod includes the previously described tobacco-containing smokable material in cut filler form.

The first filter segment is provided by gathering a 11.75 inch wide web of tobacco and carbon paper available as (P-144-BAC) from Kimberly-Clark Corp. using the filter rod forming apparatus described in Example 1 of U.S. Patent No. 4,807,809 to Pryor et al. The plug wrap for the filter segment is available as Reference No. 5831 from Ecusta Corp. The first filter segment is positioned adjacent the smokable rod.

The second filter segment is provided by gathering a 11.75 inch wide web of non-woven polypropylene web available as PP-100-F from Kimberly-Clark Corp. using the filter rod forming apparatus described in Example 1 of U.S. Patent No. 4,807,809 to Pryor et al. The plug wrap for the filter segment is available as Reference No. 5831 from Ecusta Corp. The second filter segment is positioned adjacent the first filter segment, at the extreme mouth end of the cigarette.

The cigarette outer paper wrapper exhibits an air permeability of about 0 CORESTA unit. The cigarette paper is available as P-2831-60-1 from Kimberly-Clark Corp.

The inner wrapper of the smokable rod is a tobacco-containing paper available as P-2831-23-3 from Kimberly-Clark Corp.

Smokable cigarette rods are provided using known techniques. In particular, the smokable material is circumscribed by a single layer of paper wrap. The weight of the smokable material within each cigarette rod is about 1 g.

The cigarettes are employed by burning the smokable rod such that the smokable material within the paper wrapper burns to yield smoke. When employed, such cigarettes yield very low levels of visible sidestream smoke and essentially no sidestream odor. The cigarettes do not self-extinguish during the smolder period experienced during FTC smoking conditions.

## EXAMPLE 2

Particles of calcium carbonate agglomerated with an alginate are provided as follows:

Into a blender is charged about 750 ml tap water, and then about 20 g glycerin. While the mixture is gently agitated, about 10 g of the ammo-

nium alginate described in Example 1 is slowly added thereto, so as to disperse the alginate in the water. The resulting mixture is transferred into a 1 liter jar, sealed, and gently rolled overnight to hydrate the alginate.

A slurry of 250 g precipitated calcium carbonate available as Code No. 2A from Pfizer Inc. in 250 g tap water is provided. Then, the slurry is added to 200 g of the water/glycerin/alginate mixture. The resulting slurry is agitated gently so as to provide a slurry having a smooth texture.

The slurry is cast onto a high density polyethylene sheet at a thickness of about 0.04 inch, and air dried to provide pieces of dried sheet about 6 inches by about 6 inches in size. The resulting dried sheet is hand ground to a fine particle size and screened to -50 US Mesh.

Particles of calcined agglomerated calcium carbonate are provided as follows:

Into a low shear mixer are charged about 832 parts tap water at ambient temperature, about 757 parts precipitated particulate calcium carbonate available as Albacar 5970 from Pfizer Inc., and about 267 parts molasses. The calcium carbonate has a rosettic structure and an average particle size (i.e., diameter) of about 2 microns. The molasses is available as Refiner's Syrup from Savannah Sugar Co., and has a solids/water content of about 3.7:1. The resulting mixture is agitated for about 5 to about 10 minutes to provide a slurry having a solids content of about 52 percent and a viscosity of about 1,200 cps as measured by a Brookfield LVT viscometer with cylindrical LV spindle No. 4.

The slurry is spray dried by continuously pumping the slurry at about 6 lbs./min. at a feed pressure of about 475 to about 500 psig to a spray dryer. The spray dryer is a Bowen Type commercial unit equipped with an SD-046 nozzle, and operated in a commercial mode. The inlet temperature is about 470° F, and the outlet temperature is about 260° F. The resulting spray dried particles have a generally spherical shape, and a moisture content of below about 2 percent. The particles are screened to a particle size of -70/+200 US Mesh.

The spray dried particles are placed on a 12 inch by 36 inch steel tray to a thickness of about 0.5 inch. The tray then is passed into a continuous belt furnace at a rate of about 8 to about 12 inches/min., and is subjected to heating under nitrogen atmosphere at above about 600° C for about 10 minutes, and at above about 400° C for about 20 minutes. The oven is set at about 720° C, and the tray is subjected to a maximum air temperature of about 670° C during that time. The heated particles are removed from the furnace into a cooling zone for about 1 hour under nitrogen atmosphere, and cooled to ambient temperature.

The calcined particles so collected are black,

are spherical in shape, are free flowing, and resist wetting. The particles are about 93 percent calcium carbonate, and exhibit a bulk density of about 0.5 g/cm<sup>3</sup>. The particles each are an agglomerated matrix of a plurality of precipitated calcium carbonate particles spaced within a carbonaceous material.

A tobacco-containing smokable filler material is prepared as follows:

Into a high shear blender is charged about 225 ml tap water, and into the water is dispersed about 5 g of the ammonium alginate described in Example 1. The resulting mixture is gently agitated at ambient temperature for about 15 minutes, until the alginate is hydrated. Then, about 20 g glycerin is added to the mixture, followed by about 25 ml tap water. To the mixture is added about 16.7 g of an "American blend" of tobacco cut filler which has been ground to a powder. Then, about 25 ml tap water is added to the mixture. The resulting mixture is agitated until a smooth slurry results. To the slurry is added a mixture of about 13.8 g of the calcined agglomerated calcium carbonate and about 13.8 g calcium carbonate agglomerated with ammonium alginate. The resulting slurry is agitated until the slurry exhibits a smooth texture. The resulting slurry is cast onto a high density polyethylene sheet at a thickness of about 0.025 inch and air dried.

### EXAMPLE 3

#### A. Preparation of an Agglomerated Matrix Filler

Into about 500 parts tap water at ambient temperature and maintained at high shear in a blender is charged about 5 parts ammonium alginate available as Amoloid HV. The mixture is agitated until the alginate is fully hydrated. The mixture is then transferred to an egg beater type mixer. Then, about 94.5 parts precipitated calcium carbonate, as described in Example 1, is charged into the mixture. Agitation is continued until the mixture achieves a smooth consistency. Then about 0.5 parts sodium dodecyl sulfate, from Aldrich Chemical Co., is charged into the mix. Agitation is continued for about 15 minutes at which time the mixture foams.

The slurry is cast to about a 0.05 inch thickness onto a high density polyethylene sheet and air dried. The resulting sheet is white and very brittle. It is ground and sieved to obtain a -30/+100 US Mesh particle size fraction. This material does not lose its structure when emersed in water. This is because the binder has become water insoluble. The material has a tap density, as measured with a

graduated cylinder, of about 0.26 g/cm<sup>3</sup>. The tap density of the original calcium carbonate, measured by the same method, is about 0.46 g/cm<sup>3</sup>.

#### B. Preparation of a Smokable Filler Material

Into about 200 parts tap water at ambient temperature and maintained at high shear in a blender is charged about 3.33 parts ammonium alginate as Amoloid HV and then about 4.5 parts glycerin. The mixture is agitated for about 15 minutes until the components are fully hydrated. The mixture is then transferred to an egg beater type mixer and agitated. Separately, about 60 parts of the above described agglomerated calcium carbonate is mixed with about 400 parts tap water and the resulting slurry is added to the mix. The mix is agitated gently until a smooth consistency is reached. The mix is then cast to about 0.05 inch thickness onto a high density polyethylene sheet. The resulting sheet is white and flexible. The resulting density of the sheet is about 0.39 g/cm<sup>3</sup>. The density of a comparable sheet made with the original non-agglomerated precipitated calcium carbonate is about 0.96 g/cm<sup>3</sup>. This smokable filler material then is shredded into strips about 1/32 inch wide to provide a smokable cut filler.

### EXAMPLE 4

An agglomerated matrix filler is provided as follows:

Into about 20 parts tap water at ambient temperature and maintained at high shear in a blender is charged about 5 parts HM Sugarles Pectin, Type LM-20 AS-Z, available from Hercules Chemicals. The mixture is agitated until the pectin is fully hydrated. The mixture is then transferred to an egg beater type mixer. Separately about 60 parts of precipitated calcium carbonate, as described in Example 1, is added to about 200 parts tap water. To this is added about 0.3 parts polyoxyethylene sorbitan mono-oleate available as Tween 80 from Sigma Chemical Co. is charged into the mixture. This slurry is then charged into the pectin mixture and agitated for about 15 minutes during which time the mixture foams.

The foamed slurry is cast to about a 0.05 inch thickness onto a high density polyethylene sheet and air dried. After drying the sheet, it is sprayed with a 5 percent aqueous solution of calcium chloride such that the resulting sheet now contains about 1 part calcium chloride. The resulting sheet is white and very brittle. It is ground and sieved to obtain a -30/+100 US Mesh particle size fraction. The material has a tap density, as measured with a



graduated cylinder, of about 0.28 g/cm<sup>3</sup>. The tap density of the original calcium carbonate, measured by the same method, is about 0.46 g/cm<sup>3</sup>.

The resulting agglomerated matrix filler is used to manufacture a smokable filler material, in much the same manner as described in Example 1.

## Claims

### 1. A cigarette comprising:

- (a) smokable filler material including an intimate mixture of (i) agglomerated matrix filler having an inorganic component and an organic component, and (ii) tobacco; and
- (b) a wrapping material circumscribing the smokable filler material.

2. The cigarette of Claim 1 wherein the smokable filler material includes a binding agent.

3. The cigarette of Claim 2 wherein the binding agent includes ammonium alginate.

4. The cigarette of Claim 1, 2 or 3 wherein the tobacco has the form of a tobacco extract.

5. The cigarette of Claim 4 wherein the smokable filler material includes an aerosol forming material.

6. The cigarette of Claim 1 wherein the wrapping material is a paper having a porosity of less than about 5 CORESTA units.

7. The cigarette of Claim 1 or 6 wherein the agglomerated matrix filler comprises a calcium carbonate component and an alginate component.

8. The cigarette of Claim 1 wherein the agglomerated matrix filler comprises about 10 to about 99 weight parts of a calcium carbonate component for each weight part of organic component.

### 9. A cigarette comprising:

- (a) smokable filler material including agglomerated matrix filler having an inorganic filler component and an organic component; and
- (b) a wrapping material circumscribing the smokable filler material.

10. The cigarette of Claim 9 wherein the smokable filler material includes a binding agent.

11. The cigarette of Claim 10 wherein the binding agent includes ammonium alginate.

12. The cigarette of Claim 9 wherein the wrapping material is a paper having a porosity of less than about 5 CORESTA units.

13. The cigarette of Claim 9 or 12 wherein the agglomerated matrix filler comprises a calcium carbonate component and an alginate component.

14. The cigarette of Claim 9 wherein the agglomerated matrix filler comprises about 10 to about 99 weight parts of a calcium carbonate component for each weight part of organic component.

15. A smokable filler material comprising an agglomerated matrix filler material having an inorganic component and an organic component.

16. The smokable filler material of Claim 15 including a binding agent.

17. The smokable filler material of Claim 15 or 16 including an aerosol forming material.

18. The smokable filler material of Claim 15 or 16 including tobacco.

19. The smokable filler material of Claim 15 wherein the inorganic component includes particles of calcium carbonate.

20. The smokable filler material of Claim 15 or 19 wherein the organic component includes an alginate.

21. The smokable filler material of Claim 18 wherein the tobacco has the form of an extract.

22. The smokable filler material of Claim 16 wherein the binding agent includes an alginate.

23. The smokable filler material of Claim 20 including diammonium hydrogen orthophosphate.

24. The smokable filler material of Claim 20 including residual foaming agent.

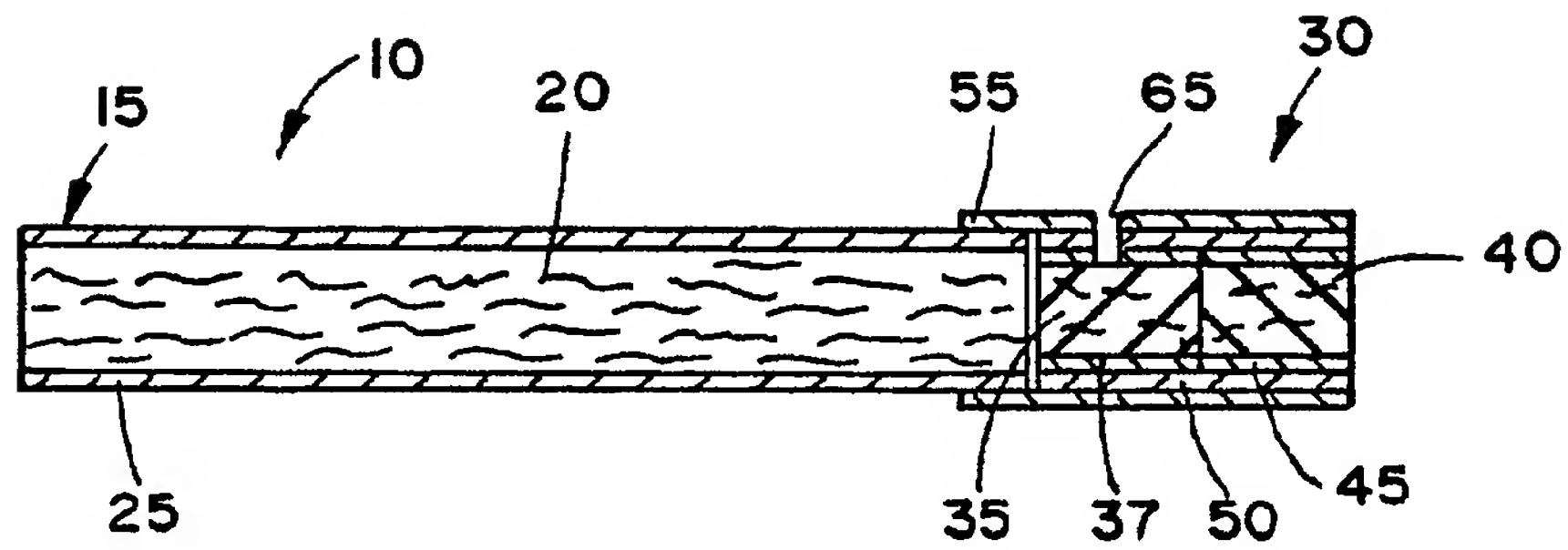


FIG. 1

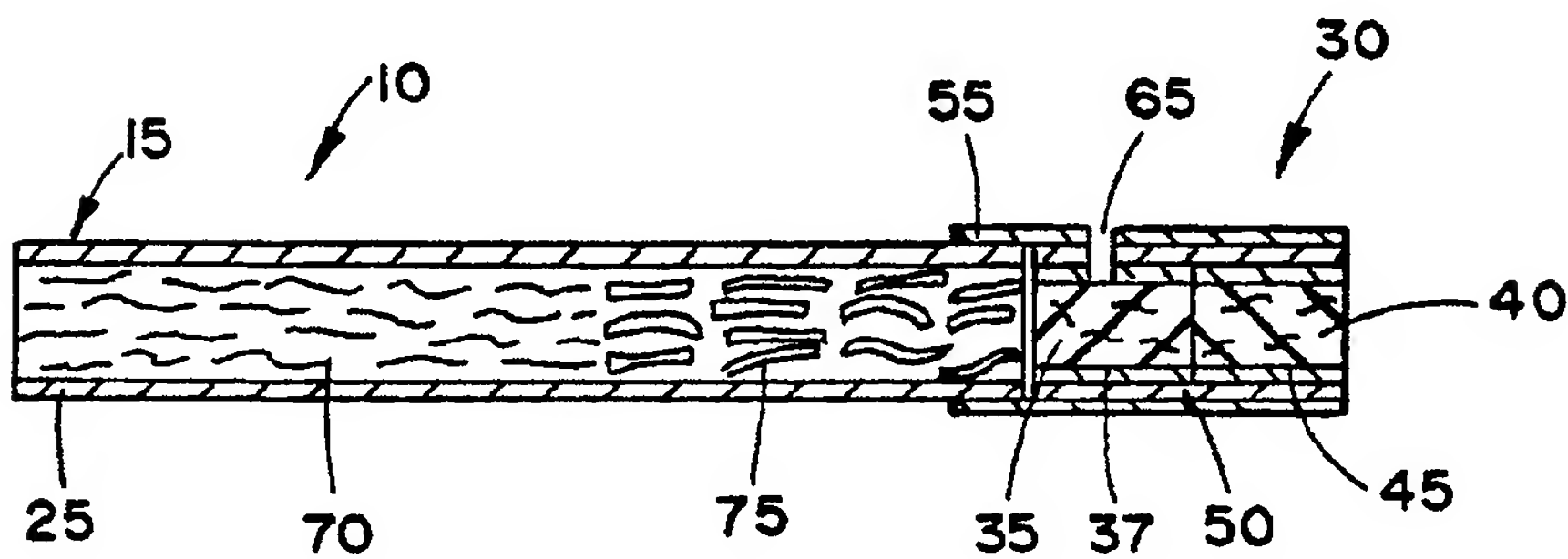


FIG. 2

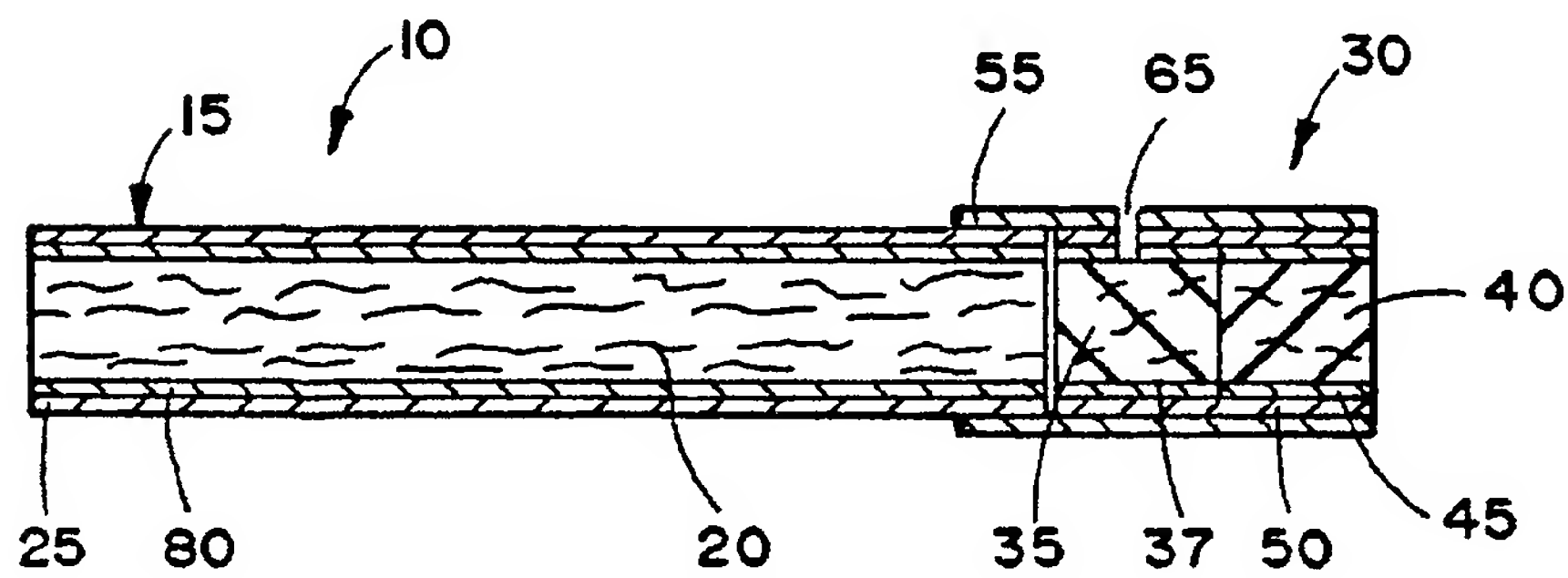


FIG. 3

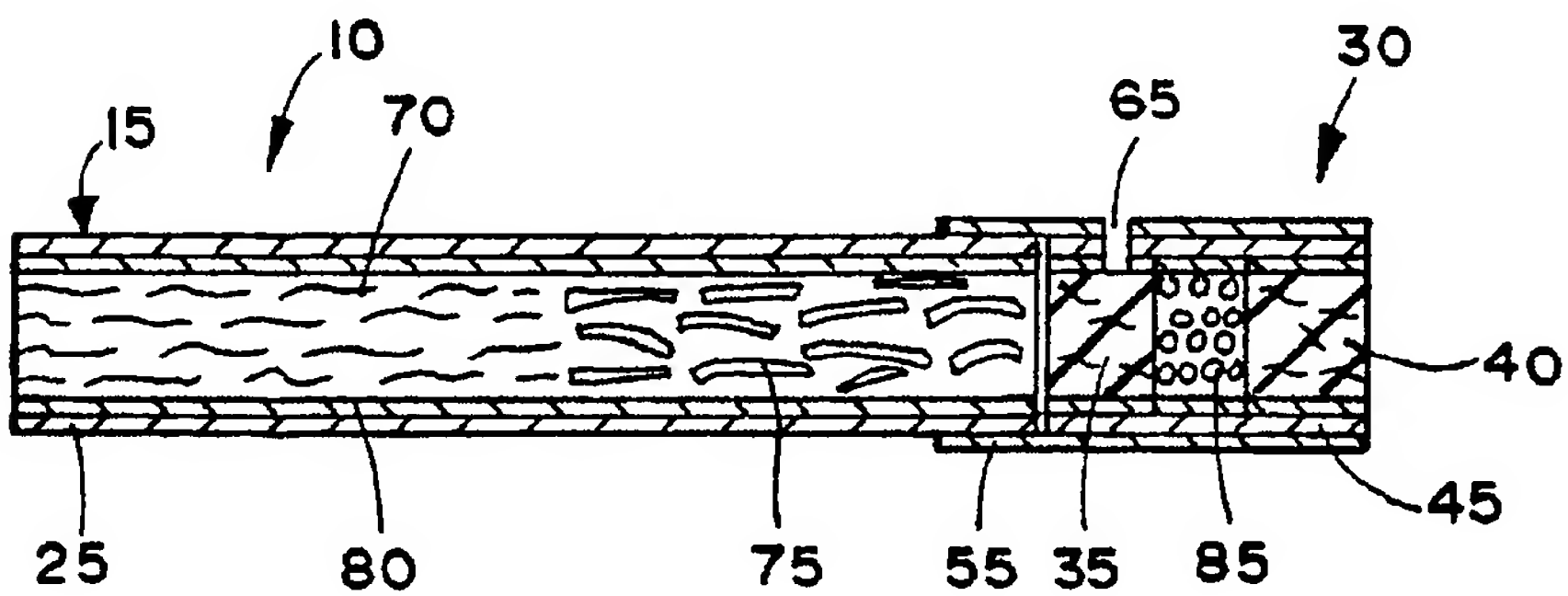


FIG. 4

Patent Application (2)  
[Stamp: 2000 yen] No symbols below  
March 26, 1973

To: Yukio MITAKE, Commissioner,  
Japan Patent Office

1. Title of Invention  
Production method of sheet tobacco

2. Inventors  
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4. List of Appended Documents  
(1) Specification 1  
(2) Copy of Application 1

Note: Documents regarding the above should be  
sent to the address below.

Patent Dept., Asahi Kasei Kogyo Co., Ltd.  
1-12 Yurakucho, Chiyoda-ku, Tokyo 100

Formal  
Examination



48 033482

## Specification

1. Title of the Invention  
Production method of sheet tobacco

### 2. Claims

A production method of sheet tobacco having a bubbly structure characterized in that a starting solution made up of water-soluble alginic acid salt, calcium carbonate, tobacco powder or substitute tobacco powder or a mixture of tobacco powder and substitute tobacco, and water is molded into a sheet, which is put in contact with an acid stronger than carboxylic acid.

. Detailed Description of the Invention

(19) Japan Patent Office  
Unexamined Patent Application Publication  
(11) Unexamined Patent Application Publication  
No.: S49-124299  
(43) Publication Date: November 28, 1974  
(21) Application No.: S48-33482  
(22) Filing Date: March 26, 1973  
Examination requested: No

(4 pages total)

JPO File No.	(52) Japanese classification:
6422 49	38 A9
6422 49	38 A0

The present invention relates to a method of molding tobacco powder or substitute tobacco powder into a sheet, characterized in that when it is molded into a sheet, a bubbly structure is brought about at the same time.

Normally, tobacco powder is produced by making powder from tobacco scraps which are generated in large quantity when cigarettes are produced, and low-grade tobacco powder is not particularly useful other than in agriculture. For this reason, since the past, attempts have been made to recycle it as tobacco by adding a caking agent and making it into a sheet.

Also, since the harmful effects of tobacco have come to light in recent years, it has become necessary to mold artificially-produced harmless



tobacco into sheets, because it is generally most often a powder.

In the past, sheets molded from tobacco powder or substitute tobacco powder (called “sheet tobacco” hereinafter) had the drawbacks of having a dense structure and being rigid and brittle. It is possible to vary the properties of sheet tobacco by changing the production method, but these drawbacks are particularly pronounced in press rolling, which is the method used most often today. Other methods have these drawbacks as well, and all methods can be said to have the problem of high density. For this reason, cigarettes made by chopping up conventional sheet tobacco are heavy in weight, and therefore have the drawback of being overpacked. They also have the drawback that when burned, the flow of air is poor due to the dense structure and combustion tends to be incomplete, and therefore the amount of nicotine or tar produced per unit length when burned is higher.

As a result of various studies on whether it is possible to provide a smoking material which is harmless to humans and has appropriate bulk and combustion characteristics without the drawbacks of the past, the inventors successfully created sheet tobacco having a bubbly structure, and thereby resolved these problems all at once.

Since the past, a widely-used method in tobacco production has been to expand and provide bulk to flattened leaf tobacco whose density has been increased. This improves combustibility and lightens flavor, and also has the effect of increasing the apparent amount of tobacco. However, with regard to sheet tobacco, technical efforts have mainly been focused only on creating sheets similar to natural leaf tobacco, and there has been no effort to provide bulk to sheet tobacco. Also, when the conventional method of providing bulk used in natural leaf tobacco is applied to the current sheet tobacco, there is the drawback that physical strength deteriorates markedly, and when subjected to conventional cigarette manufacturing machinery, it ends up being crushed into tiny pieces. Also, in the conventional method of providing bulk, sheet tobacco was temporarily produced and

then bulk was provided, but in the present invention, there is the major advantage that formation of sheet tobacco and providing of bulk – that is, manifestation of a bubbly structure – are performed simultaneously.

The first part is a substitute tobacco production method comprising the steps of making cellulose into alkali cellulose, and then heating until the pyranose ring disappears. A temperature of 160°C or above is normally required to make the pyranose ring disappear. Disappearance of the pyranose ring can be easily confirmed by the infrared absorption spectrum. For the cellulose which has lost its pyranose ring, the remaining alkali is neutralized and removed by washing with water, and when dried, it results in substitute tobacco. This substitute tobacco exhibits a dark brown color and has the form of small lumps or powder. Thus, in order to be used, it must be pulverized, the particle size must be made uniform, and it must be molded into sheets. Another example of a substitute tobacco production method which has been successfully performed is making cellulose into alkali cellulose, and then performing alkali oxidation decomposition in an oxygen-containing atmosphere until the average degree of polymerization is 95 or less, and then neutralizing this cellulose having an average degree of polymerization of 95 or less, washing it with water and drying, thereby obtaining substitute tobacco. This substitute tobacco consists of short white fibers and is not smoked as is as a substitute for tobacco, and therefore it must be molded into sheets. The present invention is extremely effective in such sheet molding of substitute tobacco.

The present invention is described in detail below.

The present invention is characterized in that a starting solution made up of water-soluble alginic acid salt, calcium carbonate, tobacco powder or substitute tobacco powder or a mixture of tobacco powder and substitute tobacco, and water is molded into a sheet, which is put in contact with an acid stronger than carboxylic acid, and it is acceptable to use substitute tobacco or a mixture of tobacco

powder and substitute tobacco instead of tobacco powder.

Here, examples of the water-soluble alginic acid salt include sodium alginate, ammonium alginate, triethanolamine alginate, magnesium alginate and the like, but otherwise, anything can be used provided that it is a salt of alginic acid that dissolves in water at room temperature. Also, an acid stronger than carboxylic acid means an acid which reacts with calcium carbonate to form calcium and a salt. Examples include strong mineral acids such as sulfuric acid, nitric acid and hydrochloric acid, and organic acids such as oxalic acid, acetic acid, citric acid, malic acid, succinic acid, tartaric acid, lactic acid and pyruvic acid. For example, when mixing the water-soluble alginic acid salt, calcium carbonate and tobacco powder, calcium carbonate may be mixed in a proportion from 10 wt% to 150 wt% of the water-soluble alginic acid salt, and it is preferred that the water-soluble alginic acid salt is used in an aqueous solution from 1 wt% to 10 wt%. The tobacco powder is added and this is made into a slurry, and creation of the starting solution is thereby complete. It is preferred that the tobacco powder is added in a quantity no more than 5 times the weight of the water-soluble alginic acid salt. If it is 5 times or more, the amount of alginic acid salt is relatively small and it does not sufficiently serve as a caking agent, and therefore the strength of the obtained sheet tobacco is diminished.

This starting solution is, for example, passed through a slit to form a sheet, and then immersed in this state into an acid stronger than carboxylic acid. The concentration in the starting solution need only be high enough to allow for sufficient penetration. In the case of a strong mineral acid, 0.1–10% is suitable, and in the case of an organic acid, 5–30% is suitable. Although sheet tobacco having a bubbly structure is obtained, acid remains attached near the surface. Therefore it must be sufficiently washed, and then dried by ordinary methods. The sheet tobacco is made up of calcium alginate and tobacco powder, and has a structure in which countless gas bubbles exist in a

continuous phase, where each bubble has substantially the same diameter.

The principle behind this is not yet understood, but the inventors hypothesize that acid penetrates into the starting solution and decomposes the calcium carbonate, thereby generating calcium ions, and then these calcium ions bond with the alginic acid to form calcium alginate which coagulates, while the carboxylic acid root which has undergone decomposition by the acid generates carboxylic acid gas, producing a bubbly structure. Therefore, the calcium ions supplied when calcium carbonate is decomposed are normally sufficient to coagulate the alginic acid, but, if necessary, calcium chloride, for example, may be additionally mixed into the acid solution, thereby supplying calcium ions from the outside, for the purpose of aiding in the coagulation of alginic acid. Also, a lubricant such as glycerol or a small quantity of inorganic reagent may be added to the starting solution.

Tobacco powder was described above, but substitute tobacco shall now be described. There are extremely few known substitute tobaccos produced by conventional techniques. The following are examples describing the substitute tobaccos produced as a result of the research of the inventors.

As described above, a requirement in the sheet tobacco having a bubbly structure of the present invention is that a water-soluble alginic acid salt is reacted with calcium ions, producing calcium alginate. Calcium alginate is a substance which is insoluble in water or acid, it gels at the same time it is formed, and the gas generated at that time is enclosed in the gel. That is, to form the sheet tobacco of the present invention, a water-soluble alginic acid salt must be used, but a caking agent such as sodium carboxymethyl cellulose of the like cannot be used.

The present invention is described in further detail below by means of examples.

#### Example 1

85 parts of water was put in a household mixer, and while it was stirred vigorously at 6000 r.p.m., 6 parts of sodium alginate and 2 parts of sedimentary calcium carbonate fine powder (equivalent to 33 wt% with respect to sodium

alginate) and 4 parts of powdered tobacco were gradually added to it a small amount at a time. In addition, 3 parts of glycerol was added, and a brown viscous suspended starting solution was obtained. This starting solution was passed through a slit 0.7 mm wide and expanded on glass plates, and each glass plate was immersed for 3 minutes in 1% hydrochloric acid aqueous solution. This was washed with water for 10 minutes and then dried for 5 hours at 110°C, and sheet tobacco having many tiny bubbles was obtained. Its void ratio was 30%.

Here, the void ratio is calculated as

$$\text{Void ratio (\%)} = \{P/(P+F)\} \times 100$$

when the area of the empty portion is taken as P and the area of the non-empty portion is taken as F in a cross-sectional photograph of the bubbly calcium alginate structure at 20x magnification. However, the values of P and F are the average values obtained from three cross-sectional photographs taken independently from the same sample.

The sheet tobacco obtained in this way was cut into 1-mm widths and rolled in ordinary rolling paper. When it was compared with conventional sheet tobacco which did not have a bubbly structure by means of a smoking test, all 20 participants in the test responded that the weight of the cigarette was lighter and the flavor was more mild.

#### Example 2

In a slurry tank equipped with a stirrer, 1 kg of wood pulp used for rayon of average degree of polymerization 636 and  $\alpha$ -cellulose content 93.7% was stirred for 20 minutes in a 3.8% slurry in a 17.8% caustic soda aqueous solution at 50°C. After that, it was squeezed in a squeeze apparatus having a vacuum filter and a press roller, after which it was continuously pulverized by a carbonate wire pulverizer, and 25 kg of alkali cellulose was obtained. This alkali cellulose had a composition with a cellulose concentration of 32.7% and a total alkali concentration of 15.6%. After that, the alkali cellulose was exposed to a 42% oxygen atmosphere at 40°C for 190 hours, and then it was made weakly acidic using an excessive

amount of 10% hydrochloric acid. It was then fully neutralized by washing with running water for 1 hour, and then dried by ordinary methods, and approximately 850 g of white short-fiber substitute tobacco was obtained. The average degree of polymerization of this substitute tobacco was 73.

10 g of this substitute tobacco, 12 g of ammonium alginate powder and 8 g of sedimentary calcium carbonate fine powder (equivalent to 33 wt% with respect to ammonium alginate) were put in a mixer, 168 g of water was added, this was vigorously stirred and mixed at 6000 r.p.m., and a milky white starting solution was obtained. This starting solution was passed through a slit 1.0 mm wide and expanded on glass plates, and each glass plate was immersed in a solution made by mixing 10% citric acid and 1% calcium chloride in a volume ratio of 3:1. This was washed with running water for 10 minutes and then dried for 5 hours at 110°C, and sheet tobacco having many tiny bubbles was obtained. The void ratio of this sheet tobacco was 28%. This sheet tobacco was cut into 1-mm widths, and it was mixed in a proportion of 3 parts sheet tobacco to 1 part cuttings for ordinary cigarette tobacco, and rolled in ordinary rolling paper to make cigarettes. When a smoking test with 20 participants was conducted, all 20 participants responded that the flavor was extremely mild.

#### Example 3

The same operations as in example 2 were repeated, except that instead of pulp for rayon containing 93.7%  $\alpha$ -cellulose, cotton linter containing 97%  $\alpha$ -cellulose was used. As a result, 26 kg of alkali cellulose was obtained. 50 g of this alkali cellulose was taken, and after being dried for 16 hours at 70°C, it was put in a 180°C atmosphere and heated for 20 minutes, and dark brown alkali cellulose was obtained. Hydrochloric acid was added to this decomposed alkali cellulose to neutralize it, and it was then washed with water 10 times, and 10 g of substitute tobacco was obtained. The same operations as in example 2 were repeated using this substitute tobacco, and mixed cigarette tobacco was obtained. When a smoking test was



conducted with 20 participants, the flavor was said to be mild and good.

Patent applicant: Asahi Kasei Kogyo Co., Ltd.

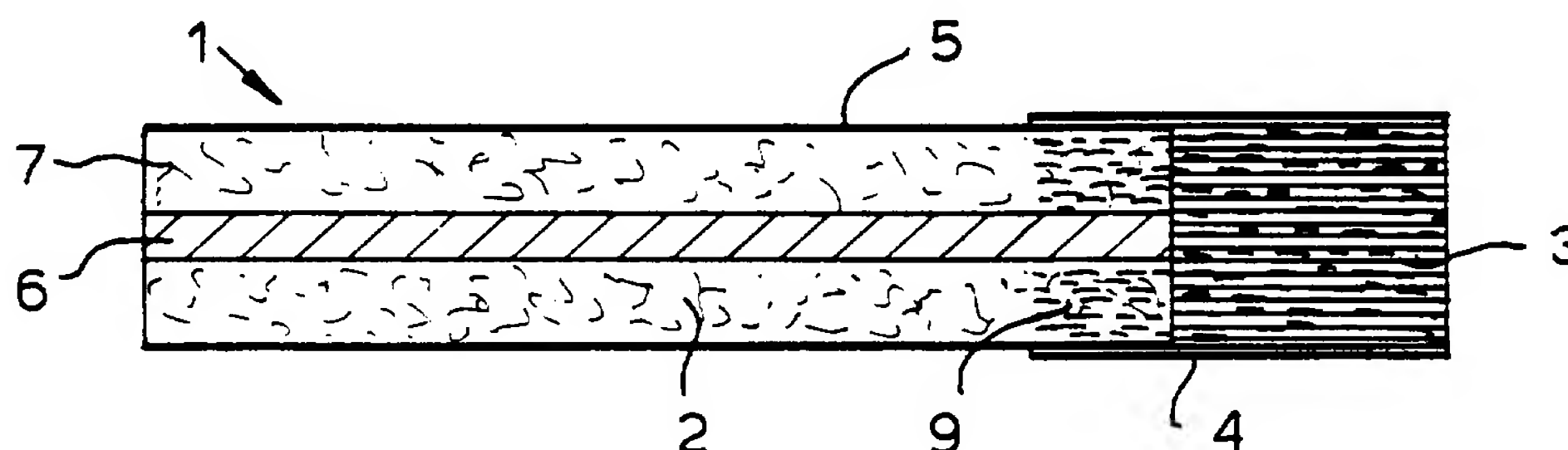
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## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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			(43) International Publication Date: 14 March 1996 (14.03.96)
(21) International Application Number: <b>PCT/GB95/02110</b>		(74) Agents: WALFORD, Margot, Ruth et al.; British-American Tobacco Company Limited, Technology Centre, Regents Park Road, Southampton SO15 8TL (GB).	
(22) International Filing Date: 6 September 1995 (06.09.95)			
(30) Priority Data: 9417970.2 7 September 1994 (07.09.94) GB 9515836.6 2 August 1995 (02.08.95) GB		(81) Designated States: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TT, UA, UG, US, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG), ARIPO patent (KE, MW, SD, SZ, UG).	
(71) Applicant (for all designated States except US): BRITISH-AMERICAN TOBACCO COMPANY LIMITED [GB/GB]; Millbank, Knowle Green, Staines, Middlesex TW18 1DY (GB).			
(72) Inventors; and			
(75) Inventors/Applicants (for US only): BEVEN, John, Lawson [GB/GB]; 1 Lambs Close, Thurleston, South Devon TQ7 3PF (GB). DITTRICH, David, John [GB/GB]; 52 Newlands Avenue, Shirley, Southampton SO15 5ES (GB). GREIG, Colin, Campbell [GB/GB]; The Old Granary, Quavey Road, Redlynch, Wiltshire SP5 2HH (GB). HOOK, Richard, Geoffrey [GB/GB]; Laurels, Palm Hall Close, Winchester, Hampshire SO23 0JL (GB). MCADAM, Kevin, Gerard [GB/GB]; Flat 1, 3 Cavendish Grove, Southampton SO17 1XE (GB). O'REILLY, Rosemary, Elizabeth [GB/GB]; 242 Bournemouth Road, Chandlers Ford, Eastleigh SO53 3HB (GB).		<b>Published</b> <i>With international search report.</i> <i>Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.</i>	
		(88) Date of publication of the international search report: 8 August 1996 (08.08.96)	

(54) Title: SMOKING ARTICLES



## (57) Abstract

The invention relates to a smoking article (1) having a high proportion of non-combustible, inorganic material and a relatively low level of visible sidestream. The smoking article comprises a substantially non-combustible wrapper (5) which extends along the full length of the smoking material rod and enwraps a combustible fuel source (6) and aerosol generating means (7) both of which extend substantially along the length of the smoking material rod. Various suitable fuel source systems and aerosol generating systems are described. The article has a visible burn line which advances along the article and produces an ash which can be removed by the smoker in the normal way.

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# INTERNATIONAL SEARCH REPORT

International Application No  
PCT/GB 95/02110

A. CLASSIFICATION OF SUBJECT MATTER  
IPC 6 A24F47/00 A24B15/16 A24D1/02

According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)  
IPC 6 A24B A24F A24D

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US,A,4 955 397 (JOHNSON ET AL.) 11 September 1990 see column 2, line 3 - line 42; figures 2,3	1,3,6, 10,13
Y	---	2,4,6-9
X	DE,A,40 09 689 (BROWN & WILLIAMSON TOBACCO CORP.) 4 October 1990 see column 3, line 46 - line 51; claim 1; figures	1,3,13
Y	---	2
	US,A,5 060 667 (STRUBEL) 29 October 1991 cited in the application see claim 1; figures	
	---	
	-/--	

☒ Further documents are listed in the continuation of box C.

☒ Patent family members are listed in annex.

### \* Special categories of cited documents :

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- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
- "&" document member of the same patent family

Date of the actual completion of the international search

6 June 1996

Date of mailing of the international search report

13.06.96

Name and mailing address of the ISA

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# INTERNATIONAL SEARCH REPORT

Int. Application No  
PCT/GB 95/02110

## C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	EP,A,0 419 975 (R.J. REYNOLDS TOBACCO COMPANY) 3 April 1991 see column 8, line 1 - line 41; claims; figures	4,6-9
X	see column 7, line 20 - column 14, line 58	43, 45-48, 50-57, 73-81, 83,84
X	--- GB,A,2 229 349 (BROWN & WILLIAMSON TOBACCO CORPORATION) 26 September 1990 see page 4, line 21 - page 5, line 9	59-64, 66-69,72
X	--- US,A,2 998 012 (LAMM) 29 August 1961  see column 2, line 31 - column 3, line 22; claims; figures	20, 23-27, 29-35, 37-40
X	--- EP,A,0 419 733 (R.J. REYNOLDS TOBACCO COMPANY) 3 April 1991  see column 7, line 32 - column 9, line 32; claims; examples 7,8 -----	43, 45-48, 50-57, 73-78, 80,81, 83,84

# INTERNATIONAL SEARCH REPORT

International application No.

PCT/GB95/02110

## Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☐ Claims Nos.:  
because they relate to subject matter not required to be searched by this Authority, namely:
2. ☐ Claims Nos.:  
because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. ☐ Claims Nos.:  
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

## Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see continuation sheets

1. ☒ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. ☐ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. ☐ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. ☐ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

- ☒ The additional search fees were accompanied by the applicant's protest.
- ☐ No protest accompanied the payment of additional search fees.



## FURTHER INFORMATION CONTINUED FROM PCT/ISA/210

invention 1: (claims 1-19)

**A smoking article with features A,B,C and D**

**A:** non combustible wrapper

**B:** a smoking material rod

**C:** combustible fuel source extending along the whole length of the smoking material

**D:** an aerosol generating means extending along the whole length of the smoking material

Invention 2: (claims 20-42 and 86-95)

**A non combustible wrapper A' with features A<sub>1</sub>, A<sub>2</sub>, A<sub>3</sub>, A<sub>4</sub>**

**A<sub>1</sub>:** inorganic filler

**A<sub>2</sub>:** binder

**A<sub>3</sub>:** plasticizer

**A<sub>4</sub>:** optional cellulosic fibre

The common concept linking 1 and 2 is a non-combustible wrapper. Even if unity could exist between 1 and 2, the common concept is not novel, therefore there is lack of unity a posteriori.

Invention 3: (claims 43-58)

**A smoking article fuel source extending along the whole length of the smoking material C' with features C<sub>1</sub>,C<sub>2</sub>,C<sub>3</sub>**

**C<sub>1</sub>:** carbonaceous material

**C<sub>2</sub>:** inorganic binder

**C<sub>3</sub>:** optional burn promoter

There is no unity between 2 and 3. The common concept between 1 and 3 is not novel (fuel source extending along the whole length of a smoking article). see eg. US-A-4955397

Invention 4: (claims 59-72)

**A smoking article aerosol generating means D' with features D<sub>1</sub>,D<sub>2</sub>,D<sub>3</sub>**

**D<sub>1</sub>:** inorganic filler

## FURTHER INFORMATION CONTINUED FROM PCT/ISA/210

**D<sub>2</sub>:** aerosol forming means

**D<sub>3</sub>:** binder

There is neither unity between 2 and 4 nor between 3 and 4. The common concept between 1 and 4, (aerosol generating means) is not novel. see eg. US-A-5060667

Invention 5: (claims 73-85)

**A smoking article aerosol generating fuel source D'' with features D<sub>1</sub>, D<sub>2</sub>, D<sub>3</sub> and D<sub>4</sub>**

**D<sub>1</sub>:** inorganic filler

**D<sub>2</sub>:** aerosol forming means

**D<sub>3</sub>:** binder

**D<sub>4</sub>:** carbon

There is neither unity between 2 and 5 nor between 3 and 5. The common concept between 1 and 5, (aerosol generating means) is not novel. see eg. US-A-5060667.

The common concept between 4 and 5 is not novel (D<sub>1</sub> - D<sub>3</sub>) see eg. EP-A-419975.

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International Application No

PCT/GB 95/02110

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